15.76.3





NCBI Conserved Domain Summary

(33)

Docket No.: PF-0346-2 CPA USSN: 09/405,940

Exhibit. No. A

RPS-BLAST 2.2.4 [Aug-26-2002]

Query= gi|1100182|gb|AAA82682.1| T-cell receptor beta [Homo sapiens]

(311 letters)

Database: oasis_sap.v1.58

4540 PSSMs; 885,521 total columns

Click on boxes for multiple alignments



Show Domain Relatives

.. This CD alignment includes 3D structure. To display structure, download <u>Cn3D</u>!

PSSMs producing significant alignments:

Score E (bits) value

● gnl|CDD|3898 smart00407, IGc1, Immunoglobulin C-Type;

77.6 6e-16

● gnl|CDD|7359 smart00409, IG, Immunoglobulin;

44.0 7e-06

● gnl|CDD|3897 smart00406, IGv, Immunoglobulin V-Type;

40.8 7e-05

gnl|CDD|7427 pfam00047, ig, Immunoglobulin domain. Members of the immunoglo...

39.1 2e-04

• gnl|CDD|3898, smart00407, IGc1, Immunoglobulin C-Type;

CD-Length = 75 residues, 98.7% aligned Score = 77.6 bits (191), Expect = 6e-16

KATLVCLATGFYPDHVELSWWVNGKEVHSGVSTDPOPLKEOPALNDSRYCLSSRLRVSAT Query: 158

PATLVCLVTGFYPPDITVTWLKNGQEVTSGVSTTDPLKDK-----DGTYFLSSYLTVSAS

Query: 218 FWQNPRNHFRCQVQFYGLSE 237

Sbjct:

1

Sbjct: 56 TW-ESGDVYTCQVTHEGLTE

• gnl|CDD|7359, smart00409, IG, Immunoglobulin;

CD-Length = 86 residues, 100.0% aligned Score = 44.0 bits (103), Expect = 7e-06

PRHKITKRGONVTFRCDPIS-EHNRLYWYRQTLGQGPEFLTYFQNEAQLEKSRLLSDRFS 85 Query: 27

Sbjct: 1 PPSVTVKEGESVTLSCEASGNPPPEVTWYKQ----GGKLLAY------SGRFS 43

Query: 86 AERPKGSFSTLEIQRTEQGDSAMYLCASSPGTSYEQYFGPGTRLTVT 132

VSRSGGNS-TLTISNVTPEDSGTYTCAATN--SSGSASS-GTTLTVL 86 Sbjct: 44

gnl|CDD|3897, smart00406, IGv, Immunoglobulin V-Type;

CD-Length = 80 residues, only 77.5% aligned Score = 40.8 bits (95), Expect = 7e-05

Query: 51 LYWYRQTLGQGPEFLTYFQNEAQLEKSRLLSERF&AEE--PKGSFSTLEIQRTEQGDSAM 100 Sbjct: 18 VSWVRQPPGKGLEWLGYIGSDVSYSEA-SYKGRVTISKDNSKNDV-SLTISNLRVEDTGT 75

Query: 109 YLCA 112 Sbjct: 76 YYCA 79

• gnl|CDD|7427, pfam00047, ig, Immunoglobulin domain. Members of the immunoglobulin superfamily are found in hundreds of proteins of different functions. Examples include antibodies, the giant muscle kinase titin and receptor tyrosine kinases. Immunoglobulin-like domains may be involved in protein-protein and protein-ligand interactions. The Pfam alignments do not include the first and last strand of the immunoglobulin-like domain.

CD-Length = 66 residues, 98.5% aligned Score = 39.1 bits (90), Expect = 2e-04

Query: 35 GQNVTFRC--DPISEHNRLYWYRQTLGQGPEFLTYFQNEAQLEKSRLLSDRFSAERPKGS 92 Sbjct: 1 GESVTLTCSVSGYPPDPTVTWLRNGKELELGVLVTS------SSSGRTSTS 45

Query: 93 FSTLEIQRTEQGDSAMYLCA 112 Sbjct: 46 SLSLTISSVTPEDSGTYTCV 65

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